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Difference In-Gel Electrophoresis: A High-Resolution Protein Biomarker Research Tool

David S. Gibson, David Bramwell, and Caitriona Scaife

Summary

Difference in-gel electrophoresis (DIGE) is a recent adaptation of conventional two-dimensional gel electrophoresis (2-DE) that incorporates novel fluorescent labels, has multiplex attributes, and boasts software-assisted image analysis. Combined, these characteristics offer significant benefits in accuracy and reproducibility to quantify differential protein expression levels between biological samples. The DIGE technique and materials required to perform it are described in detail within. The principles behind consistent gel image acquisition and reliable image analysis are also considered. Within the context of biomarker and drug target discovery, this method simplifies analysis, increases sample throughput, and represents a reliable 2-DE platform.

Key Words: Biomarker; Cy dye; DIGE; fluorescent difference in-gel electrophoresis; proteomics; two-dimensional gel electrophoresis

1. INTRODUCTION TO DIFFERENCE IN-GEL ELECTROPHORESIS

Two-dimensional gel electrophoresis (2-DE) is an established platform that facilitates the analysis of complex protein mixtures. O'Farrell was first to introduce high-resolution two-dimensional electrophoresis by resolving proteins to individual isoelectric point and molecular weight coordinates (*1*). The main asset of this method is that it provides a global view of the state of proteins within a sample. In theory, thousands of proteins can be visualized at once, giving a unique qualitative “map” or “fingerprint” of changes

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01 between given samples. Though many developments, such as standardized
02 immobilized pH gradients, have led to vast improvement in inter-run
03 consistency, deficiencies in sensitivity and spot matching have necessi-
04 tated further adaptation using fluorescent stains. Comparison between large
05 groups of conventionally (silver or Coomassie) stained gels is complicated
06 by spot to spot warping, caused by variations in sodium dodecyl sulfate-
07 polyacrylamide gel electrophoresis (SDS-PAGE) gel casting, electric and
08 pH fields, and thermal fluctuations during electrophoresis. This leads to
09 problems in spot matching and necessitates multiple gel replicates to prevent
10 assumptions on mismatched proteins. In other words, gel to gel hetero-
11 geneity makes it difficult to distinguish with confidence between variations
12 in the technique and those of genuine induced biological change, such as
13 in disease states (2). Difference in-gel electrophoresis (DIGE) addresses a
14 number of these issues in that two to three samples can be subjected to
15 exactly the same running conditions within a single gel. Unlu *et al.* developed
16 DIGE to allow a more direct and reproducible comparison between protein
17 samples, differentiated by prelabeling with spectrally resolvable fluorescent
18 cyanine, or Cy, dyes (3). The Cy dyes are charge matched with the residues
19 they bind to within the proteins of a given sample and have similar
20 molecular weights (0.5 kDa), thus result in only slight gel shifts. The Cy
21 dyes are based on extended organic ring structures and hence are highly
22 hydrophobic. Concerns with protein precipitation prior to electrophoresis
23 have been surmounted by using a “minimal labeling” strategy, whereby
24 binding is limited to only 1% to 2% of lysine residues available within a
25 sample (4).

26 Excitation of each fluor allows the creation of a digital image of each
27 individually labeled sample. These dyes give additional validity to the two-
28 dimensional technique in the form of higher sensitivity, wider dynamic
29 range, and linearity of detection. Detection limits of 0.025 ng are possible,
30 with a dynamic range around five orders of magnitude. One of the strongest
31 features of the technique, however, is the ability to include an internal pooled
32 standard, which is loaded on all gels within an experiment (5). The internal
33 standard permits the linking of all gels in an experiment, thus offering
34 more reliable and intuitive software-assisted comparisons. The accuracy of
35 protein quantification between samples is increased dramatically, and much
36 smaller changes in protein expression can be studied with greater confi-
37 dence. Evaluations of DIGE alongside traditional and more recent proteomic
38 methods using isotope-coded or isobaric tags (cICAT and iTRAQ) reveal
39 that it remains competitive in sensitivity and can be used with confidence
40 as a platform for drug discovery and development (4,6).

2. DIGE EXPERIMENT CONSIDERATIONS

Because three cyanine dyes are available, up to three separate protein samples can be labeled per gel. A pairwise analysis and organization could also be used (akin to gene chip analysis), where control versus drug-treated samples are labelled with Cy3 and Cy5 only. When normalization of expression levels is desired across a number of different experiments and within the one experiment, adequate quantities of each sample should be available to create a common pooled internal standard. The internal standard can be distinguished from experimental samples by labeling with Cy2 dye. Anomalies in spot intensity due to preferential labeling can be eliminated by randomized or reciprocal labeling, in which half of each experimental group is labeled with Cy3 and the other with Cy5 (7). In order to distinguish intrinsic, interindividual biological variation from genuine changes in protein expression, biological replicates should be included in each experimental group. A recent study focused on the DIGE technique has shown that a minimum of four replicate gels is required to maintain a 95% chance of avoiding false negatives, when a twofold change in expression is considered significant (8).

3. SAMPLE PREPARATION

Plasma and synovial fluid are used in this chapter to illustrate and describe the steps required for the purification and minimal fluorescent labeling of body fluid samples. For details of how to prepare cell lysates, with both minimal and saturation types of labeling, one can refer to the Ettan DIGE system user manual (9). The following reagents and conditions have been used in our laboratory to produce reliable data with clinical relevance to patient outcome but could also be applied to prospective drug trial to monitor therapeutic response. Sample preparation should be consistent and kept as simple as possible to reduce inter-run inconsistencies. Protein modifications during sample preparation must be prevented, particularly degradation due to endogenous proteolytic enzymes. Such changes in samples analyzed by gel-based approaches can translate into misleading artifact spots with novel molecular weights.

3.1. Sample Purification and Assay

Cellular or particulate material should be removed from the body fluid by centrifugation prior to any further purification steps. This circumvents contamination by sub-proteomes other than that of the body fluid that is to be analyzed. Endogenous protease activity should be inactivated for reasons already eluded to above. A number of approaches are possible, with

01 varying consequences to the resulting sample integrity. Protease inhibitor
 02 cocktails (such as the Complete Protease Inhibitor Cocktail Tablets; Roche) AQ1
 03 can be used to inactivate a wide variety of degradative enzyme classes
 04 including cysteine, matrix metallo and serine proteases. This remains our
 05 preferred method of body fluid stabilization, and an adaptation is now
 06 also available, in the form of a blood tube with proprietary inhibitors for
 07 immediate and convenient sample protection (BD P100). AQ2 Some authors,
 08 however, caution against their use in certain applications, as artifacts can
 09 result from modified protein charge, or peptide-based inhibitors such as
 10 leupeptin may interfere with mass spectrometry analysis (*10,11*). Proteases
 11 may also be inactivated by high or low pH extremes with Tris buffer or
 12 trichloroacetic acid (TCA), respectively, or alternatively total protein can
 13 be precipitated by TCA/acetone. In balance though, protein yield may be
 14 diminished by incomplete precipitation or resolubilization. Once stabilized,
 15 salts can be removed from protein samples (if higher than 10 mM) by
 16 dialysis with low-molecular-weight cut-off membranes, though if analysis
 17 of small peptides is desired, precipitation could be implemented. Other
 18 macromolecules such as lipids, polysaccharides, and nucleic acids should be
 19 removed by organic solvent, unless present at low concentrations (as with
 20 plasma). The sample can be lyophilized if concentration is necessary (5 to
 21 10 mg/mL is an ideal protein concentration, though labeling of 1 mg/mL is
 22 possible) and resuspended in a minimal quantity of DIGE-compatible lysis
 23 buffer [DLB; 30 mM Tris, 7 M urea, 2 M thiourea, 4% (w/v) CHAPS, pH
 24 8.5]. Ampholytes and dithiothreitol (DTT) are omitted from the lysis buffer
 25 prior to the labeling reaction as both primary amines and thiol groups will
 26 compete with the proteins for the available Cy dye. The pH of the sample
 27 to be labeled is also critical to the reaction, so check that the sample pH is
 28 8.5 by spotting on a pH indicator strip and, if necessary, make drop-wise
 29 adjustments with dilute sodium hydroxide. The concentration of protein in
 30 each sample should be assayed either by Bradford reagent or using the
 31 proprietary Ettan 2D Quant kit (GE Healthcare). AQ3

32 **3.2. Sample Labeling**

34 Aside from the pH and protein concentration recommendations already
 35 made, the efficiency of minimal dye labeling is dependent on the ratio of
 36 dye to protein (400 pmol Cy dye to 50 µg protein is cited in the DIGE
 37 user manual; GE Healthcare). The Cy dye fluors should be reconstituted
 38 in anhydrous dimethyl formamide under the manufacturer's guidelines to
 39 create a 1 mM stock solution. Each has a characteristic deep color as follows:
 40 Cy3, red; Cy5, blue; and Cy2, yellow (as shown in **Fig. 1**). The sample and
 41 dye quantities required for a six-gel, three-dye pilot experiment are shown

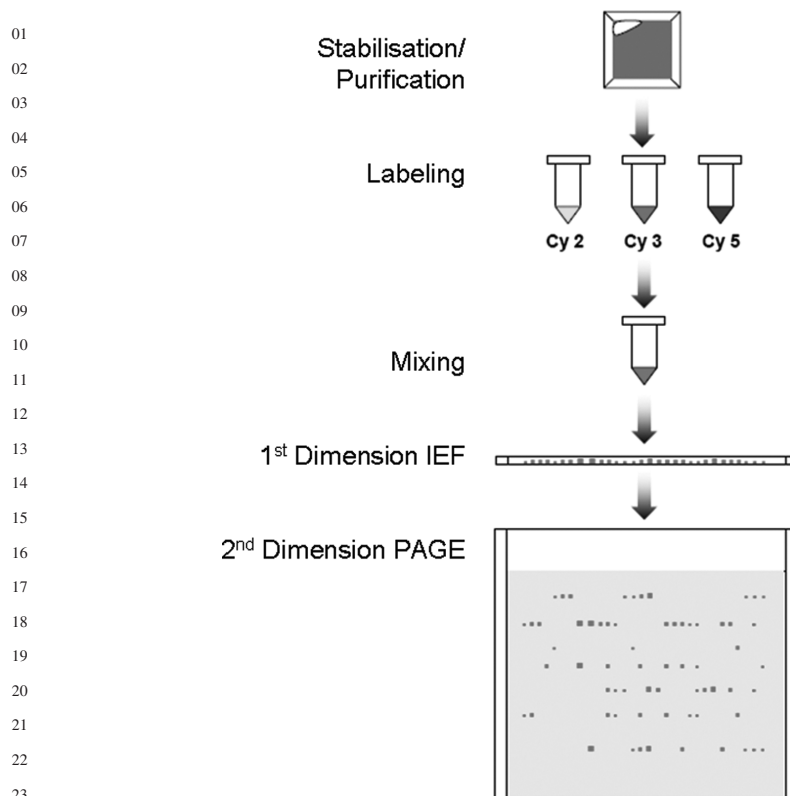


Fig. 1. Schematic representation of the laboratory procedures involved in a typical DIGE experiment.

in the following worked example (**Table 1**). Paired plasma and synovial fluid samples from six patients (A–F) are labeled with Cy3 and Cy5. An experimental design incorporating randomization of sample labeling and loading across gels is demonstrated to avoid systematic errors. An internal pooled standard is generated by combining equal amounts of all matched plasma and synovial fluid samples, followed by Cy2 dye labeling. Sufficient pooled internal standard is prepared to allow enough aliquots for each gel in the experiment. It is also prudent to create a slight excess (10% to 20%) of each dye reaction to ensure a complete aliquot is loaded on each gel. Thus for the individual plasma and samples, 60 μg is labeled with Cy3 or Cy5, but only 50 μg will be loaded of each. A single internal standard is therefore prepared, which comprises 30 μg of each of the 12 samples (6 plasma and 6 synovial fluid) and labeled with Cy2 dye. Before labeling, it is recommended that all sample concentrations are normalized to 10 $\mu\text{g}/\mu\text{L}$, to make subsequent pipetting easier (**Table 2**).

Table 1

The Sample and Dye Quantities Required for a Six-Gel, Three-Dye Pilot Experiment Are Shown for a Three-Dye Run Analyzing Plasma (PL) and Synovial Fluid (SF) from Six Patients (Anonymized as A, B, C, D, E, and F)

Gel	Cy2 pooled standard	Cy5	Cy3
1	50 µg (4.17 µg of each sample SF A–F and PL A–F)	50 µg of PL C	50 µg of SF D
2	50 µg (4.17 µg of each sample SF A–F and PL A–F)	50 µg of SF A	50 µg of PL F
3	50 µg (4.17 µg of each sample SF A–F and PL A–F)	50 µg of PL B	50 µg of SF E
4	50 µg (4.17 µg of each sample SF A–F and PL A–F)	50 µg of SF C	50 µg of PL A
5	50 µg (4.17 µg of each sample SF A–F and PL A–F)	50 µg of PL D	50 µg of SF B
6	50 µg (4.17 µg of each sample SF A–F and PL A–F)	50 µg of SF F	50 µg of PL E

Beforehand: Label Reaction Tubes

1. Label 12 Microfuge tubes (0.5 mL) as **Table 2** (*SF A–F, PL A–F*) for preparation of samples prior to labeling.
2. Label a second fresh set of 12 Microfuge tubes as above, for the labeling reaction.
3. Label one Microfuge tube as *PS* for the pooled standard.

Sample Preparation

4. Aliquot volumes of each sample equivalent to 100 µg into each of the first set of *SF* or *PL* individually labeled tubes. Adjust protein concentrations of all samples to 10 µg/µL by addition of DIGE compatible lysis buffer (DLB), as shown in **Table 1**.
5. Aliquot 6 µL of each normalized sample (60 µg) from the above, into each of the second fresh set of *SF* or *PL* individually labeled tubes and 3 µL (30 µg) of each sample into the one *PS* labeled tube. This gives a total of 36 µL (or 360 µg protein) in the pooled standard tube (*PS*).

Sample Labeling

6. The Cy dyes are diluted from 1 mM stock concentration to a working concentration of 400 pmol/µL with DMF, and 1.2 µL of the Cy3 and Cy5 dyes is added to individual samples in a randomized fashion as described in **Table 1**. An aliquot of 7.2 µL of 400 pmol/µL Cy2 is added to the pooled standard tube. (*Note*: Only reconstitute minimal quantities of Cy dye working dilutions for

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Table 2
The Volumes of Samples and Diluent Required for Individual and Pooled Dye Reactions Are Shown*

Sample	[Protein] $\mu\text{g}/\mu\text{L}$	Volume required for 100 μg (μL)	Volume of DLB for 10 $\mu\text{g}/\mu\text{L}$ (μL)	Volume add to PS labeling tube (μL)	Volume add to individual labeling tubes (μL)	Labeling tube
SFA	21.0	4.8	5.2	3	6	SF A
SF B	29.2	3.4	6.6	3	6	SF B
SF C	17.6	5.7	3.3	3	6	SF C
SF D	11.8	8.5	1.5	3	6	SF D
SF E	17.4	5.7	4.3	3	6	SF E
SF F	27.6	3.6	6.4	3	6	SF F
PL A	39.4	2.5	7.5	3	6	PL A
PL B	35.5	2.8	7.2	3	6	PL B
PL C	23.2	4.3	5.7	3	6	PL C
PL D	26.5	3.8	6.2	3	6	PL D
PL E	23.6	4.2	5.8	3	6	PL E
PL F	31.4	3.2	6.8	3	6	PL F
Total for pooled internal standard				36	PS	

*Sample concentrations are normalized to 10 $\mu\text{g}/\mu\text{L}$.

the experiment as any remaining stock can be kept for future work at -20°C for 3 months.)

7. All labeling reaction tubes should be mixed thoroughly by pipette and vortex and pulse-centrifuged to collect the mixture at the bottom. Incubate the labeling reaction tubes on ice in the dark for 30 min. (*Note:* Subsequent exposure of all dye reactions to ambient light, whether in IPG strip or gel, should be minimized to prevent degradation/bleaching of the fluorophore.)
8. Add 1.2 μL of 10 mM lysine to each of the Cy3 and Cy5 dye reactions and 7.2 μL to the Cy2 dye reaction to stop the labeling. Again, mix and centrifuge briefly before incubating for a further 10 min on ice in the dark. The labeling reaction is now complete, and labeled samples can be stored for up to 3 months at -70°C in a light protected container, if not used immediately.

4. FIRST-DIMENSION ISOELECTRIC FOCUSING (IEF)

The ampholytes and DTT that had been omitted prior to the labeling are added at this point in the form of a 2X sample buffer. The sample is denatured with dithiothreitol (DTT) and a volume equivalent to 50 μg of each individual Cy3 and Cy5 labeled sample are then combined with 50 μg of the Cy2 pooled internal standard. The mixture is subsequently rehydrated onto 24-cm immobilized pH gradient (IPG) strips for highest resolution (sample in-gel rehydration). Samples with larger quantities of high molecular weight proteins, alkaline proteins, or hydrophobic proteins are likely to be poorly absorbed into the IPG strip gel matrix and would benefit substantially from cup loading detailed elsewhere (*12*).

Beforehand: Prepare Buffers

1. Prepare 2X sample buffer [8 M urea, 130 mM DTT, 4% (w/v) CHAPS, 2% (v/v) IEF ampholytes 4–7] and rehydration buffer (8 M urea, 13 mM DTT, 4% (w/v) CHAPS, 1% (v/v) IEF ampholytes 4–7).
2. Remove pH 4–7 IPG strips from freezer to thaw on bench and ensure rehydration tray is level. Note strip numbers and label six fresh microcentrifuge tubes as **Table 1 (I–6)**.

Pooling Samples, Strip Rehydration, and Isoelectric Focusing

1. Add equal volumes of 2X sample buffer to individual Cy3 and Cy5 labeled samples (8.4 μL each) and to the Cy2 labeled pooled standard (50.4 μL), mix and leave on ice for 10 min (each tube now has 50 μg labeled protein in 14 μL).
2. Aliquot 14 μL of each Cy3, Cy5, and Cy2 labeled samples to be focused on the same IPG strip into the tubes as indicated in **Table 1**. Add 408 μL of rehydration buffer to each tube, mix, and centrifuge briefly.
3. Pipette each mixture into a separate channel of the rehydration tray. Peel off the protective cover from the IPG strip and carefully lower it gel side down into the rehydration buffer-sample; remove any air bubbles with a pipette tip.

Table 3
Isoelectric Focusing Conditions Appropriate to Proteins Soluble Within the Acidic Range pH 4–7 for 24-cm IPG Strips

Voltage mode	Voltage(V)	Duration (h:min)	Volt-hours (kVh)
1. Step	3500	—	75
2. Gradient	1000	0:10	—
3. Step	8000	1:00	—
4. Step and hold	100	>24 h	—

4. Overlay each strip with ~2 mL of IPG strip cover fluid to prevent evaporation and slide on the plastic cover. Cover with aluminum foil and incubate overnight (or at least 10 h) at room temperature. Low voltage applied during rehydration can improve entry of high molecular weight protein (**13**).
5. After adequate rehydration, use clean forceps to remove the IPG strips from the rehydration tray. Blot off excess fluid and carefully position the IPG strips (gel side up) within the ceramic manifold of the IPGphor isoelectric focusing unit (GE Healthcare). Ensure that the positive (anodic) end of each strip is oriented toward the anode of unit.
6. Apply one filter paper electrode pad, which has been moistened with de-ionized water (remove excess fluid with blotting paper), to each end of the IPG strip in such a way as to overlap the gel by approximately 3 to 5 mm. Place the respective electrodes over the filter paper pads and clip firmly in place and overlay each strip with cover fluid to prevent dehydration (108 mL for whole manifold).
7. Close the IPGphor IEF unit safety lid and the instrument running conditions can be programmed. We suggest conditions appropriate to proteins soluble within the acidic range pH 4–7, for 24-cm IPG strips (**Table 3**). Current should be limited to 50 μ A per IPG strip. Other run conditions for the various strip sizes and pH ranges are available [Ettan IPGphor manifold user manual (80-6499-52 edition AO); GE Healthcare].
8. Once the strips have been focused, they can be equilibrated straight away for the second dimension or stored for several months at -80°C between plastic sheets to prevent damage to the brittle frozen strips.

5. SECOND-DIMENSION SDS-PAGE (2-DE)

In order to improve protein migration from focused IPG strip to the second-dimension separation gel, it is important to equilibrate the strips in a buffer containing sodium dodecyl sulfate (SDS), urea, and glycerol. In the first equilibration step, DTT is added to completely reduce any remaining disulfide bonds, whereas iodoacetamide is added in the second step to

alkylate the resultant sulfhydryl groups thereby preventing reoxidation, which may complicate downstream mass spectrometry identification. In addition, the iodoacetamide “mops up” any free DTT, which may otherwise cause point-streaking artifacts, apparent in silver-stained gels. The Protean Plus Dodeca multiple gel unit (Biorad) is used in the worked example, as it can accommodate up to 12 large-format slab gels, has an in-built buffer recirculation and cooling system and plate electrodes for consistent resolution. A recipe for a 12% homogenous polyacrylamide Tris-glycine gel is also given using stabilized, high-purity Protogel reagents (National Diagnostics), though gradient gels and other buffer systems may give better resolution for select protein molecular weight ranges (14,15). It should be noted that the nature of the Cy dyes used in the DIGE technique necessitate the use of specialized low-fluorescence glass plates. The glass plates require thorough cleaning with dilute alcohol and de-ionized water using lint-free tissues to remove any dried salts or gel fragments prior to casting.

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Beforehand: Prepare Equilibration Buffer and Gels

1. Prepare the stock SDS Equilibration buffer [50 mM Tris pH 8.8, 6 M urea, 30% (v/v) glycerol, 2% SDS (w/v)] (10 mL per strip will be required, excess can be aliquoted and stored at -20°C for future use).
2. Prepare the working-strength Gel Running buffer [25 mM Tris base, 192 mM glycine, 0.1% SDS (w/v)] (up to 25 L is required to fill the Dodeca gel tank).
3. Prepare 600 mL of gel casting solution (sufficient for six 1.0-mm, 12% gels; 100 mL per gel) [240 mL 30% (w/v) acrylamide/methylene bisacrylamide solution (37.5:1 ratio), 156 mL 4X Laemmli Resolving Gel buffer (0.375 Tris-HCl pH 8.8, 0.1% SDS final concentration, 197.4 mL de-ionized water, 240 μL TEMED, 2.4 mL of 10% (w/v) ammonium persulfate (APS)]. Pass the solution through a 0.2- μm filter and degas under vacuum, prior to the addition of APS.
4. Apply waterproof adhesive tape to the sides of assembled glass plates and spacers, place into the gel caster with plastic sheet spacers, and gradually pour in the prepared mixture. Carefully layer ~ 2 mL of water-saturated butanol on top of each gel to remove bubbles and create a level surface. Flush the butanol off the gels with double de-ionized water after approximately 1 h and cover the casting unit with tin foil. Ideally, gels should be cast the day before use to ensure complete polymerization.
5. Prepare a 1.0% agarose solution in Gel Running buffer with 50 mg bromophenol blue incorporated per 100 mL. Dissolve the agarose by a short incubation in a microwave on low-medium power.

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IPG Strip Equilibration

1. Prepare Equilibration buffer A by dissolving 100 mg of DTT per 10 mL of stock SDS Equilibration buffer (5 mL needed per IPG strip). Dispense a 5-mL

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01 aliquot of buffer A into equilibration tubes and place the IPG strips carefully
02 into each. (Disposable plastic 10-mL pipettes with the conical tip broken off
03 are sufficiently long; reseal with parafilm between incubations.) Incubate the
04 strips at room temperature with gentle rocking for 15 min, then decant the
05 buffer.

- 06 2. Prepare buffer B by dissolving 250 mg of iodoacetamide per 10 mL of stock
07 SDS Equilibration buffer (5 mL needed per IPG strip). Dispense a 5-mL aliquot
08 of buffer B into each equilibration tube and reseal. Equilibrate the strips for a
09 further 15 min at room temperature with gentle rocking, decant the buffer, and
10 proceed to electrophoresis section.

SDS-PAGE

- 11
12 1. Rinse the equilibrated IPG strips in Gel Running buffer and using forceps place
13 each strip across the top of a gel, such that the plastic backing of the strip
14 makes contact with the back glass plate and the anodic end of the strip is at the
15 top left of the gel. A thin spatula can then be used to maneuver the strip down
16 into the well, with care so no bubbles are introduced between strip and gel.
17 A good tip is to push one end of the IPG strip down, between the glass plates,
18 so it makes contact with the top of the gel and gradually push the opposite end
19 down so the strip sits level on the surface.
- 20 2. Layer approximately 2 mL of premelted 1% agarose sealing solution carefully
21 across the strip, again try to eliminate any air bubbles trapped between the strip
22 and gel. Allow to cool and solidify for 5 min and repeat the procedure for the
23 outstanding IPG strips.
- 24 3. Insert the prepared gel cassettes into position in the electrophoresis unit (this
25 can be made easier by prewetting each cassette by dipping it briefly into the
26 Gel Running buffer). Ensure the buffer chamber is filled to the manufacturer's
27 recommended level and place the safety lid on paying attention to the orientation
28 of the electrodes.
- 29 4. Start the electrophoresis with 2 W per gel for 45 min followed by 17 W per
30 gel for 4 h, both at 20°C. Alternately, for convenience, the gels can be run
31 overnight at 0.75 W per gel. (Such an overnight run can take anything from 18
32 to 21 h.) The run should be terminated when the bromophenol blue dye front
33 reaches the bottom of the gel.
- 34 5. In preparation for scanning the gels, the scanner instrument should be switched
35 on and left to warm up for 30 min and the glass plate should be thoroughly
36 cleaned with lint-free tissues. As DIGE gels are scanned while they are in the
37 glass cassettes, these too should be cleaned carefully to remove any residual
38 running buffer, gel smears, and so forth, from the surface of the plates. If
39 not scanned immediately, gels can be individually wrapped in cling film and
40 stored at 4°C but should be imaged as soon as possible to avoid signal loss and
41 dissipation of focused spots. Gels stored at 4°C should also be allowed to come
up to room temperature before scanning to avoid any condensation forming on
the surface of the glass.

6. IMAGE ACQUISITION

The capture of gel image data is a critical stage in the whole analysis workflow that can often present a source of easily prevented experimental noise. It is possible to make careless choices at this stage that can result in a 200-fold drop in sensitivity. A variety of DIGE-compatible gel scanning instruments based on charge-coupled device (CCD) or photomultiplier tube (PMT) detection technologies are available (GE Healthcare Typhoon; Syngene, ChemiGenius; Fuji, FLA 1500; Biorad, FX Molecular Imager). Depending on the make of scanner, there are several parameters to optimize; the spatial resolution, the dynamic range, the scan content, and background offsets. These features are therefore briefly discussed and recommendations made in each case to improve the process of image acquisition.

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6.1. Spatial Resolution

Image (or spatial) resolution relates to the number of pixels (picture elements) displayed per unit length of a digital image and is often measured in dots per inch (dpi) or in micrometers (the size of the area each pixel represents). Images with a higher spatial resolution are composed of a greater number of pixels and have more image detail than those of lower spatial resolution. It is important to be aware that variations in spatial resolution will not only affect the final appearance of the image but will also impinge on the quality of spot detection and the accuracy of any subsequent quantitative measurements.

At low resolutions, there are fewer pixels available to represent each spot, and as a result, spot detection and quantitative accuracy will be compromised. Image (spatial) resolution is illustrated in more detail in **Fig. 2**. Higher resolution means that more pixels, and hence more data, are available for the analysis: the spot highlighted in red in **Fig. 3** is represented by 63 pixels at a relatively low 100 dpi resolution (**Fig. 2A**), compared with 485 pixels at a higher 300 dpi resolution (**Fig. 2B**).

There is, however, a maximum resolution, which once exceeded produces minimal additional information. Once resolution is sufficient to adequately represent the smallest features, any further increases in spatial resolution simply increase the ability to represent the system “noise.” In addition, every doubling in spatial resolution quadruples the amount of data that has to be processed, which can cause problems in processing speed and file and memory management. For example, a typical 20 × 20 cm DIGE single dye image captured at 100 dpi versus 300 dpi would result in diverse file sizes of 1.2 Mb and 10.6 Mb, respectively (both at 16-bit depth).

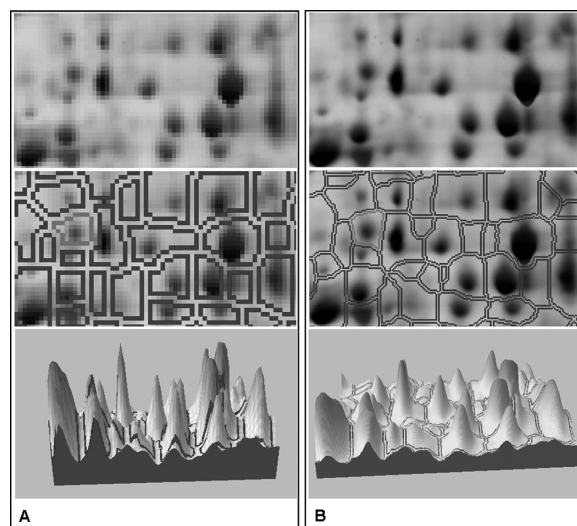


Fig. 2. Portions of two-dimensional gel image scans showing typical image quality, spot outline, and three-dimensional spot intensity (top to bottom) at 100 dpi (8-bit) and 300 dpi (16-bit) resolutions.

To summarize, in most situations, 300 dpi or 100 μm will provide an image that is large enough for accurate analysis and small enough for efficient processing. However, if your gels are small (e.g., minigels), then you may need to increase the resolution to achieve this. As a rule of thumb, the active area of the gel (i.e., the area of spot material) should fall in the range 1000 to 1800 pixels in both horizontal and vertical directions. This range provides a good trade-off in information content and analysis performance.

6.2. Dynamic Range (Bit Depth)

Also referred to as color depth, bit depth or pixel depth is the number of bits used to represent the color (grayscale, intensity levels) of each pixel in an image. Greater bit depth allows a greater range of colors or shades of gray to be represented by a pixel. If possible, scan at 16-bit rather than 8-bit. The bit depth of a 16-bit image (65,536 levels of grayscale) compared with an 8-bit image (256 levels of grayscale) results in enhanced sensitivity and accuracy of quantification for less-abundant proteins. The possible grayscale levels available along with the resultant dynamic range (orders of magnitude) for the types of images commonly used in two-dimensional gel image analysis are indicated in **Table 4**. In reality, the images displayed on the computer

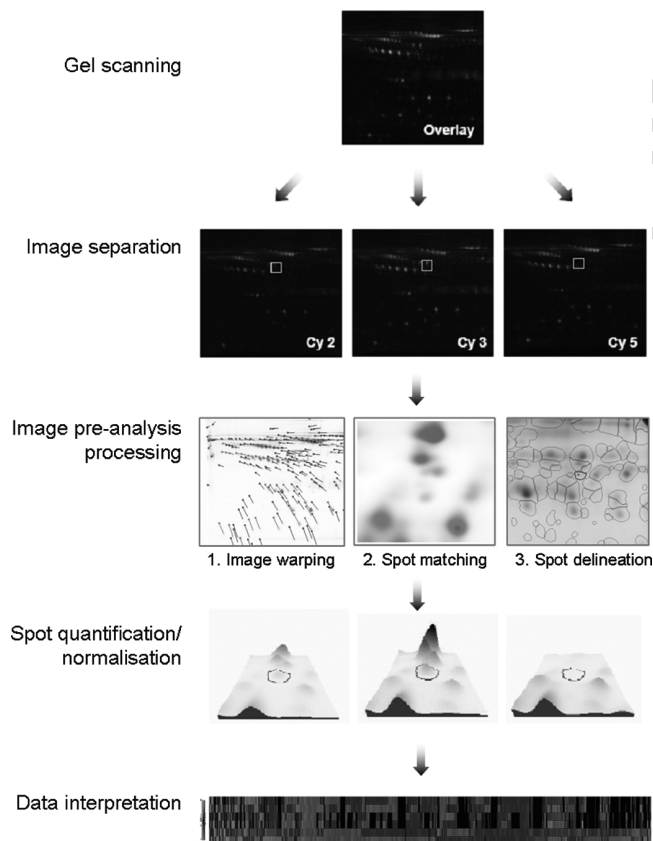


Fig. 3. Schematic representation of the processes involved in DIGE image analysis. (see Color Plate 4, following p. x)

screen will only be represented in 256 shades of gray, and so an 8-bit image will look identical to a 16-bit image by eye. However, image analysis software can distinguish between the different levels of gray. As a rule, the more levels of gray represented in an image, the better the ability to differentiate low-abundance spots from background, and the greater the quantitative accuracy. This is further illustrated in **Fig. 2**, comparing spot detection in an identical area on the same two-dimensional gel, captured at 8-bit and 16-bit, respectively.

The dynamic range can be adjusted in CCD camera systems by altering the exposure time and in laser-based systems by fine tuning the voltage of the PMT detector. The dynamic range should be optimized to maximize the use of available grayscale values. Aim for the maximum gray levels

Table 4
The Resultant Grayscale Levels and Dynamic Range (Orders of Magnitude) Are Shown for a Range of Image Bit Depths Commonly Used to Analyze Two-dimensional Gels

Bit depth	Intensity (grayscale) levels	Orders of magnitude	Percent of 16-bit scan
8	256 (2^8)	2.4	0.39
10	1024 (2^{10})	3.01	1.56
12	4096 (2^{12})	3.6	6.25
16	65,536 (2^{16})	4.8	100

in the image to be 5% to 10% less than those available. Scanner response curves can be nonlinear, and inconsistent settings can cause issues. For DIGE experiments without a Cy2 internal standard, chose settings that optimize the dynamic range for each stain and keep these consistent. When optimizing the dynamic range, it is important to avoid saturation effects. Saturation occurs when gray levels exceed the maximum available. When a spot becomes saturated, any differences in high pixel intensities cannot be resolved, and the spot appears truncated when viewed in three dimensions. No reliable quantitative data can be generated from a saturated spot, and saturated spots may also have an overall effect on normalization if included in later analysis.

6.3. Scan Content

Wherever possible, it is best to try to keep the area scanned as valid gel area. It is common to see scans where there are lots of scanner bed, labels, and so forth, in the captured image. Some scanners automatically adjust the scan settings based on what they “see” so a significant proportion of the dynamic range is lost by representing scanner bed or labels. It may be best to switch off auto gain control features or, at least, outline the areas you are interested in so the scanner optimizes only the region of interest. Extra “non gel” areas provide no useful information and should be cropped prior to image analysis. These scanning artifacts can skew the image statistics, “steal” dynamic range, increase storage requirements, and cause extra work in manual stages of analysis. Another good tip for consistency (and to save time later) is to always scan gel images using the same orientation and with the same settings.

Postscan processing of two-dimensional gel images using Adobe Photoshop or other general image processing software should be avoided, as

these do not maintain the integrity of your original data, and any calibration information contained in the image file will be lost. The manipulations may make the images “look better” to the human eye but are simply transforming the original data. If the images look bad in the first place, then you should try to optimize the scanning not manipulate the images digitally. If possible, use GEL or IMG/INF files formats rather than generating TIFF files. The former often contain additional grayscale calibration information, which will not be included in the TIFF version. In any case, do not use JPEG file storage format as it is a standard for “lossy” image compression, which is optimized to allow the loss of information that is least noticeable to the human eye. This does not mean it does not affect measures made by computers. Lossy compression throws information away and manipulates the image data. Converting a JPEG image back to a TIFF is not a solution; once the image has been compressed in this way, the data has been lost and cannot be retrieved.

7. IMAGE ANALYSIS: SOFTWARE PRINCIPLES AND WORKFLOWS

Image analysis of conventional silver or Coomassie stained two-dimensional gels of individual samples can be highly subjective and very time consuming, due to inherent unpredictable distortions between gels. These inconsistencies prevent the perfect alignment and matching of spots between gels. On the other hand, DIGE-derived images from the same gel are precisely superimposable, and gel to gel spot matching is uniquely assisted by the internal standard (Cy2 labeled). So by virtue of the prelabeling and comigration of differentially labeled samples, variation in spot “coordinates” and intensity are accounted for, and gel images analysis is much more efficient (*16*). It is important to understand the principles that traditional software applications apply, highlight their limitations, and provide a recommended analysis workflow (depicted in **Fig. 3** and *see* Color Plate 4, following p. x).

7.1. Image Warping and Spot Matching

The biggest problem in the image analysis of a gel-based experiment is data alignment. This is easy to see by comparing the state of the art statistical analysis for array-based experiments and the sorts of analysis performed on gel-based systems. The key difference between the two workflows is that the exact locations of data points are known in the array scenario. In traditional two-dimensional gel analysis, the alignment issue is tackled by a between gel spot matching stage. Most traditional analysis strategies follow

01 these key steps; detect spots on each gel individually, attempt to match the
02 spots across the experiment (possibly with the assistance of whole image gel
03 warping), measure spots, apply some form of normalization, apply statistical
04 analysis (17). The recommended workflow (**Fig. 3**) is distinguished by
05 initially warping gels, then matching and delineating spots, and imposing
06 the same spot outlines across all gels prior to quantification.

07 The main problem with the traditional workflow is that it is not able
08 to provide data of sufficient quality required to perform advanced statis-
09 tical analysis. The core issue is focused on missing values; that is, a data
10 point (usually a spot) that is not available across all samples. A recent
11 study measured 42% missing values (i.e., not experimentally induced data
12 omission) for a 16-gel experiment (18). It has also been shown that the
13 number of missing values increases with the number of replicates (19). This
14 produces the predicament of reducing the extent and quality of data, as
15 the investigator endeavors to improve statistical power by increasing the
16 number of replicates. In the traditional analysis workflow, missing values
17 arise from two main sources: (1) the same measurements not being taken
18 from each gel (usually due to spot detection) and (2) the measurements not
19 being correctly matched between gels.

20 The same measurement not being taken from each gel has two main
21 effects. The first is down to the fact that in traditional analysis scenarios,
22 each gel has spots detected in isolation. This can lead to inconsistent results
23 because essentially the pattern is determined from a single instance and
24 as such is more prone to technical variance. The second issue is usually
25 attributed to experimental conditions where a spot has zero expression in
26 one or more of the groups. In this case, a spot will not be detected on an
27 individual gel basis, and we are left with a hole in the data. Strangely, the
28 proteomics community tends to differ with standard scientific practice with
29 zero expression spots preferring to have no measurement or “unmatched”
30 rather than measuring the value “zero.” This is analogous to measuring
31 air temperature and saying “there was no temperature” when it hits zero
32 instead of measuring and recording zero. This particular stance also forces
33 a multiple instead of a unified statistical framework approach to analysis,
34 which is laborious, prone to bias, and still may be suboptimal.

36 7.2. Spot Delineation/Outline

38 Geometric correction alone does not solve all of the issues as we would
39 still be prone to threshold of detection issues on a per gel basis and also
40 not matching spots that were not expressed in certain groups. The obvious
41 next step is to stabilize and standardize the spot detection across all of the

01 images within an experiment. The geometric correction combined with an
02 “ensemble,” experiment-wide, spot pattern would go a large way to not only
03 removing the bulk of missing values but also removing a large amount of
04 manual intervention. We would also gain most in the low-expression spots
05 that are harder to detect consistently on a per gel basis. If we apply geometric
06 correction and derive a suitable outline for spots that are not undergoing
07 experimentally induced variation, then the software can handle more than
08 half (usually more) of the spots in an experiment fully automatically. This
09 is essentially reverse logic where it is more efficient to bias the analysis to
10 discount what is not changing rather than to optimize for what may be. This
11 also removes the situation where you have to spend a lot of time editing
12 a series of spots to find “they don’t change.” Considering the efficiency
13 of overall analysis, applying the same spot outlines across geometrically
14 corrected images is advantageous, and the benefits increase as the number
15 of spots subject to experimentally induced expression changes decreases.
16 A restriction imposed by the same spot outlines is that the area considered
17 to contain a spot is consistent across images. This means that there are only
18 going to be issues if the expression change is so large that it cannot be
19 represented by the same area across the gels. For spots in isolation, this is
20 not an issue as a larger area can be used (any extra pixels included in the
21 larger boundary will be zero in the smaller spots). Tight clusters of spots
22 that vary slightly in location and potentially patterns of spots that overlap
23 when considered across all groupings may however prove problematic and
24 require additional editing. Interestingly, posttranslational modification shifts
25 of proteins are better handled by the same outlines workflow than the
26 traditional approach, as one can apply a robust statistical framework to
27 discovering them. Therefore, contrary to current practice and “gut feel,”
28 it appears to be more favorable to apply the same spot outline pattern to
29 geometrically corrected gels and robustly analyze the bulk of the spots
30 within the experiment in a “first pass” with minimal manual intervention
31 and hence bias.

32 Once the same outlines analysis has been completed, one will have a list
33 of “interesting” areas. These will either be completely satisfactory with the
34 automatic outlines or it may be deemed that editing should be considered. It
35 is advisable that the initial analysis be completed fully without intervention
36 and this analysis saved. From this, a list of areas for further examination
37 should be made and these explored in a mode that adds to the information
38 and not replaces it. At this point, we may consider editing outlines. The
39 editing process is less biased when one uses the same outline across all
40 gels within the experiment. This is because the volume of a spot is affected
41 by the number of pixels chosen to include in its boundary; if imposed

inconsistently between groups, it is easy to introduce unintentional bias. Because there is a much reduced requirement on manual intervention, one can spend a lot more time on the areas that matter. Feedback from multiple labs has shown that differential editing is rarely necessary at all and, if it is, tends not to be a major factor in the results of the experiment. Areas where this is deemed necessary should always be treated with caution and may be a target area for future experiments where ~~the~~ zoom gels are used or alterations to the sample preparation improve the data quality in these areas.

In summary, the suggested workflow is

1. Geometrically correct your gels.
2. Create a spot pattern representative of all of the spots within your experiment.
3. Analyze the spots completely.
4. Complete statistical analysis.
5. Create a copy of the experiment and edit/apply differential outlines on areas where it is deemed strictly necessary.
6. Complete statistical analysis.

8. PREPARATIVE GELS FOR SPOT PICKING

Because DIGE offers superlative sensitivity to quantify minute differences in protein expression levels, relative to conventional staining techniques, sample load is reduced accordingly (from as low as 75 μg to 200 μg per gel). Once spots of interest have been identified, it is then necessary to run a “preparative” gel with a higher sample loading (0.5 mg to 2.0 mg) ensuring sufficient protein in these spots to obtain reliable identifications by mass spectrometry (20). A spot “pick list” can be generated, such that isoelectric point and molecular weight coordinate data derived from the DIGE “analytical” gels can be transposed onto a Coomassie or silver stained gel.

9. CONCLUSION

Whereas the concept of DIGE is relatively novel, it has unique attributes that make it particularly suitable in the drug and biomarker discovery process. Because the technique is not dependent on antibody-protein or oligonucleotide-RNA/DNA affinity, it does not preclude the target identity, therefore represents changes in the sample without preemptive bias. Although extended multiplex capability and an increased dynamic range are desirable, it remains the most sensitive validated gel-based proteomic tool with direct relevance to innovation in clinical and pharmaceutical research. Improvements in dedicated analytical software have greatly increased the throughput and confidence in data derived from DIGE images. DIGE holds

01 much promise in the challenge to uncover new molecular targets, screen
 02 putative drug efficacy, and monitor therapeutic response for a wide range
 03 of debilitating diseases.

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UNCORRECTED PROOF

Chapter-9

Query No.	Page No.	Line No.	Query
AQ1	192	02	Provide location of Roche.
AQ2	192	07	Provide name and location of manufacturer.
AQ3	192	31	Provide location of manufacturer.
AQ4	194	31	We have italicized the underlined words throughout the text and table. Please check.
AQ5	197	Table 3	In Table 2, define acronym DLB.
AQ6	198	05	Provide location of manufacturer.
AQ7	198	10	Provide location of manufacturer.
AQ8	198	26	Define acronym TEMED.
AQ9	200	08	Provide locations for Syngene and Fuji.
AQ10	200	18	Explain parts A and B in the legend add part labels to Fig. 2 legend.
AQ11	208	28	In Ref. 9, provide location of manufacturer as publisher.